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Enhanced photocatalytic NOx oxidation and storage under visible-light irradiation by anchoring Fe₃O₄ nanoparticles on mesoporous graphitic carbon nitride (mpg-C₃N₄)



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ABSTRACT

Several mesoporous graphitic carbon nitride (mpg- C_3N_4) photocatalysts were synthesized by using a hard-templating method comprising thermal polycondensation of guanidine hydrochloride over silica spheres at three different temperatures (450, 500 and 550 °C). After structural characterization of these mpg- C_3N_4 photocatalysts, they were tested in NO(g) photo-oxidation under visible (VIS) light. The effects of polycondensation temperature on the structure and photocatalytic performance of mpg- C_3N_4 in NO photo-oxidation were studied. The results revealed that polycondensation temperature has a dramatic effect on the photocatalytic activity of mpg- C_3N_4 in NO photo-oxidation, where mpg- C_3N_4 synthesized at 500 °C (mpg-CN500) showed the best performance in NO_x abatement as well as a high selectivity towards solid state NO_x storage under VIS light illumination. Photocatalytic performance of the mpg-CN500 was further enhanced by the anchoring of 8.0 \pm 0.5 wt.% Fe₃O₄ ananoparticles (NPs) on it. Fe₃O₄/mpg-CN500 photocatalyst showed both high activity and high selectivity along with extended reusability without a need for a regeneration step. Enhanced photocatalytic NO_x oxidation and storage efficiency of Fe₃O₄/mpg-CN500 photocatalyst was attributed to their mesoporous structure, high surface area and slow electron-hole recombination kinetics, efficient electron-hole separation and facile electron transfer from mpg-CN500 to Fe₃O₄ domains enhancing photocatalytic O₂ reduction, while simultaneously suppressing nitrate photo-reduction and decomposition to NO₂(g).

1. Introduction

Atmospheric pollution is considered to be one of the major threats for modern society. Anthropogenic air pollutants such as nitrogen oxides (NO_x) induce ozone production in troposphere and cause acid rains. Particularly, nitric oxide (NO_x) and nitrogen dioxide (NO_x) severely affect human respiratory and immune systems [1–3]. Although NO_x emissions have been strictly regulated by the environmental protection agencies, the recommended value of $\leq 0.2\,\mathrm{ppm}$ is often exceeded in urban settings [4]. Therefore, finding improved approaches for NO_x removal from the atmosphere is essential. Although NO_x abatement can be performed efficiently using thermal catalytic technologies (*i.e.* selective catalytic reduction/SCR and NO_x storage and reduction/NSR) at elevated temperatures [5–8], an important challenge

is the abatement of gaseous NO_x species under ambient conditions (*i.e.* at room temperature and atmospheric pressure).

In this regard, semiconductor photocatalysis presents an appealing alternative, since the major requirements for these technologies are only sunlight, water and oxygen which are naturally present in the atmosphere in abundance [9]. This approach has been already implemented in advanced construction materials to combat urban NO_x pollution, using mainly titania (TiO₂) based photocatalysts [10–12]. Despite its favorable properties like chemical inertness, long-term stability and low cost, TiO₂ has a typical band gap of 3.0–3.2 eV. This large band gap enables the harvesting of mostly UV light, which constitutes only 4–5% of the incoming solar energy [13,14]. More importantly, in photocatalytic NO_x abatement applications, TiO₂ has a low selectivity towards NO_x storage in solid state and tends to oxidize NO(g) into a

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more toxic product, $NO_2(g)$ [15]. Therefore, it is necessary to develop alternative photocatalysts harvesting a broad wavelength window with high efficiency and stability. Moreover, the proposed $DeNO_x$ materials should be abundant, accessible, easy to synthesize, stable and non-toxic in order to justify their implementation on a large scale.

Among such potential photocatalytic systems, two-dimensional (2D) graphitic carbon nitride (g-C₃N₄) is one of the strong contenders owing to its metal-free nature, thermal stability, non-toxicity, low-cost and relatively narrow band gap (2.7 eV, 454 nm) falling in the visible range of the solar spectrum [16]. These unique properties make g-C₃N₄ as an attractive candidate for various applications in the field of solar energy conversion and environmental remediation [17]. The uniquely versatile 2D-layered polymeric structure of g-C₃N₄ which mainly consists of carbon and nitrogen atoms along with a minor amount of H atoms (e.g. at termination and defects sites) can be synthesized using different approaches including polycondensation, pyrolysis, solvothermal and physical/chemical vapor deposition [18,19]. However, g-C₃N₄ has a very low surface area (10-50 m²/g) which limits its practical photocatalytic applications. To overcome this drawback, mesoporous graphitic carbon nitride (mpg-C₃N₄) can be prepared by template-assisted methods that are particularly beneficial as they enable fine-tuning of the surface, electronic, and morphological properties of g-C₃N₄ and also allow the design of highly porous structures with a large surface area (150-400 m²/g) [20]. Mesoporous photocatalytic structures with a large surface area can increase the number of exposed active sites of the photocatalyst, which in turn, can improve the photocatalytic activity through enhanced solar light absorbance capacity.

It is well known that photo-oxidation and storage of NO(g) in the form of solid state nitrate species is a complex process and involves several intermediates such as nitrogen dioxide (NO₂(g)) [15]. It is important to note that NO₂(g) is far more toxic than NO(g) as it can contribute towards formation of secondary pollutants like acid rain. Thus, extensive release of NO₂(g) at the end of the photocatalytic process can nullify the overall photocatalytic NO_x abatement effect [21]. Therefore, it is of paramount importance to design photocatalysts which restrict the formation and/or emission of toxic side-products such as NO₂(g). Designing such a photocatalyst is a challenge as increased selectivity can also sacrifice photocatalytic activity.

Photocatalytic architectures containing $\mathit{mpg}\text{-}C_3N_4$ can be enhanced using a variety of synthetic strategies. Among these strategies, addition of non-precious transition metals is one of the most effective and convenient approaches to introduce desired properties into the $\mathit{mpg}\text{-}C_3N_4$ photocatalyst system. Magnetite (Fe $_3O_4$) is a common form of iron oxide containing both Fe 2 + and Fe 3 + cationic species in a spinel structure. It has been demonstrated that Fe $_3O_4$ could be utilized as a promising catalytic promoter due to its unique electronic and magnetic properties [22–24]. In this context, in the current study, we integrate Fe $_3O_4$ nanoparticles (NPs) and 2D $\mathit{mpg}\text{-}C_3N_4$ nanosheets in a single nanocomposite system as an efficient and inexpensive photocatalyst for airborne NO $_x$ abatement.

In the present study, several $mpg\text{-}G_3N_4$ photocatalysts were synthesized by using a hard-templating method comprising thermal polycondensation of guanidine hydrochloride (GndCl) over silica spheres at three different temperatures (450-550 °C). The structure of as-synthesized photocatalysts were characterized by using a multitude of analytical techniques. Thereafter, photocatalytic performances of these photocatalysts were tested in NO photo-oxidation under environmentally relevant reaction conditions. Moreover, the photocatalytic performance of the mpg-CN500 in NO oxidation was further enhanced by the assembly of 8.0 \pm 0.5 wt.% Fe₃O₄ NPs on it.

2. Materials and methods

Iron (III) acetylacetonate [Fe(acac)₃, 99.9%], oleylamine (> 70%), benzyl ether (98%), Ludox* HS-40 (40 wt.% colloidal SiO₂ suspension in water) and Titanium (IV) oxide (P25, \geq 99.5%) were purchased from

Sigma-Aldrich. Guanidine hydrochloride (GndCl, 98%) and ammonium hydrogen difluoride (NH_4HF_2 , 98.5%) were purchased from Alfa-Aesar and Fluka, respectively. All chemicals used in this work were used as received without further purification.

2.1. Synthesis of the mpg-C₃N₄

The mpg-C₃N₄ photocatalysts were prepared by the thermal polycondensation of GndCl in the presence of Ludox® HS-40 colloidal silica as a hard template. In a typical synthesis campaign [25], 4.0 g of GndCl was dissolved in 4 mL distilled water in a glass vial and added dropwise into 10 g of Ludox[®] HS40 colloidal silica under vigorous stirring. The resultant mixture was kept at 50 °C for overnight. Obtained white solid was crushed in a ceramic mortar and was placed into a quartz crucible with a cover for annealing. The crucible was heated in a horizontal quartz-tube oven at 450, 500 and 550 °C for 2 h with a ramp rate of 4-5 °C min-1 under argon flow. The resultant products were denoted as mpg-CN450, mpg-CN500 and mpg-CN550, respectively. Next, the yellow solid powders attained via this approach were reacted with 4 M, 200 mL of NH₄HF₂ (aq) solution for two days to remove the silica template. Finally, obtained powders were repeatedly washed with water and ethanol in order to remove the residual reactants and dried overnight at ca. 50 °C in a vacuum oven.

2.2. Synthesis of Fe₃O₄ nanoparticles

Monodisperse Fe_3O_4 nanoparticles were synthesized by using a well-established protocol published elsewhere [26]. In a typical synthesis, 3 mmol of $Fe(acac)_3$ was dissolved in 15 mL of benzyl ether and 15 mL of oleylamine. The resultant solution was heated to $120\,^{\circ}C$ and kept at this temperature for 1 h under nitrogen atmosphere. Next, the resultant mixture was quickly heated to $300\,^{\circ}C$, and kept at this temperature for 1 h. Then, the solution was cooled to room temperature. To purify and separate the Fe_3O_4 nanoparticles (NPs), ethanol was added to the reaction mixture and the mixture was centrifuged for $12\,\text{min}$ at $8500\,\text{rpm}$. Separated Fe_3O_4 NPs were dispersed in hexane for the further use.

2.3. Incorporation of the mpg-C₃N₄ with Fe₃O₄ NPs

In a typical procedure, 50 mg of monodisperse Fe $_3O_4$ NPs were added to 10 mL of hexane and then mixed with 150 mg of mpg-CN500 sample which was initially dispersed in 30 mL of ethanol. The resultant ethanol/hexane mixture was sonicated for 2 h to ensure adsorption of the Fe $_3O_4$ NPs onto mpg-CN500. Next, the resultant mixture was centrifuged at 7500 rpm for 10 min and the separated nanocomposite (containing both mpg-CN500 and Fe $_3O_4$) was washed with ethanol for several times and dried under vacuum. These samples will be referred as Fe $_3O_4/mpg$ -CN500 in the rest of the text. Fe $_3O_4$ content of the synthesized Fe $_3O_4/mpg$ -CN500 materials was determined by ICP-MS analysis as 8.0 ± 0.5 wt.%. This apparently small Fe $_3O_4$ loading is due to the removal of oleylamine (OAm), a long chain (C18) organic molecule which was used as surfactant for the synthesis of Fe $_3O_4$ NPs. In other words, 50 mg Fe $_3O_4$ sample actually included a much lesser amount of pure Fe $_3O_4$.

2.4. Structural characterization

Crystal structures of the synthesized materials were analyzed using a PANalytical Empyrean XRD diffractometer equipped with Cu-K α radiation (40 kV, 45 mA, $\lambda=1.54051$ Å). Attenuated Total Reflectance (ATR) Fourier transform-infrared (FTIR) spectra were collected using a Bruker Alpha Eco-ATR spectrometer in the frequency range of 400–4000 cm $^{-1}$. The optical absorption properties of the samples were investigated *via* Diffuse Reflectance UV-VIS (DR-UV-VIS) Spectroscopy using a Cary 5000 UV-VIS-NIR Spectrometer equipped with a Varian

Cary 2500 Internal Diffuse Reflectance (DR) Accessory. The Brunauer-Emmett-Teller (BET) specific surface area (SSA) measurements of the synthesized catalysts were carried out using nitrogen adsorption-desorption isotherms obtained with a Micromeritics 3Flex surface area and pore size analyzer. Prior to SSA analysis, all samples were outgassed in vacuum for 2h at 150 °C. X-ray photoelectron spectroscopy (XPS) experiments were performed with a SPECS PHOIBOS hemispherical energy analyzer. A monochromatic Al-Kα Xray excitation source (15 kV, 400 W) was employed during the XPS data acquisition. Transmission electron microscopy (TEM), Scanning transmission electron microscopy (STEM), High Angle Annular Dark Field (HAADF) imaging and Energy Dispersive X-Ray (EDX) analysis experiments were carried out at 300 kV using an FEI Technai G2 F30 Transmission Electron Microscope equipped with BF-/DF-STEM-EDX modules. The photoluminescence (PL) spectra were performed at room temperature using a Jobin-Yvon Horiba Fluorolog-3 equipped with a Hamamatsu R928 P detector and a 450 W ozone-free Osram XBO xenon arc lamp. The excitation wavelength was 390 nm which is well above the sample band gap. The fluorescence was monitored at a right angle relative to the excitation. Sonication experiments was performed in a VWR USC 900 T Ultrasonic Cleaner Bath (9.2 L, 45 kHz) at room temperature. Temperature of the sonication bath was kept below 30 °C via ice addition into the bath. Without any ice addition, the final temperature of the sonication bath reached up to 40 °C after 2 h.

2.5. Photocatalytic activity measurements

The photocatalytic DeNOx experiments were performed at room temperature in flow mode by considering the experimental requirements that were reported in the ISO 22197-1 standard [27]. Inlet gas mixture that was introduced to the reactor (Fig. S1) contained 0.750 standard liters per minute (SLM) N₂ (g) (purity: 99.99%, Linde GmbH). 0.250 SLM O₂ (purity: 99%, Linde GmbH) and 0.010 SLM NO (100 ppm NO (g) diluted in balance N2 (g), Linde GmbH). In order to obtain the gas flow values given above, mass flow controllers (MFCs, MKS1479 A for N2 (g) and O2 (g) and Teledyne HFC-202 for NO (g) diluted in N2 (g)) were utilized so that the typical total gas flow over the photocatalyst was stabilized at 1.010 SLM ± 0.05 SLM, where the NO (g) content of the inlet gas mixture was fixed at 1 ppm. The pressure inside the reactor was kept at ca.1 bar and measured via a MKS Baratron 622B capacitance manometer. Humidity of the inlet gas mixture was also carefully controlled by dosing varying amounts of water vapor into the inlet gas mixture (i.e. before the reactor entrance) with the help of a Perm Select (PDMSXA-2500) semi permeable membrane module attached to an external variable-temperature water chiller/recycler for controlling equilibrium vapor pressure of water. Typical relative humidity (RH) of the reactor was kept within 50 \pm 3% at 23 \pm 2 °C, measured at the sample position using a Hanna HI 9565 humidity analyzer. Changes in the NO, NO2, and total NOx gas concentrations at the outlet of the reactor were monitored using a chemiluminescent NO_x analyzer, Horiba Apna-370 with a 0.1 ppb sensitivity and 1 Hz (i.e. 1 measurement per second) detection speed.

For the experiments performed with VIS illumination, a 35 W metal halide lamp (HCI-TC 35 W/942 NDL PB 400–700 nm range, Osram) was utilized. Since the currently used VIS light source also emitted a limited but detectable flux of UV light, a commercial VIS-transparent UV-blocker/filtering film (LLumar window film UV CL SR PS (clear)) was placed on top of the reactor during the VIS-light experiments. This was crucial for ruling out any contribution from UV photons during the VIS-light illumination. Numerically, without using UVA filter, the UVA flux measured by the UVA probe (LP471 UVA, DeltaOhm) within 315–400 nm wavelength range was 2.092 W/m², while this value dropped by %99.93 to 0.015 W/m² in the presence of the UVA filter (with which the experiments were carried out). The incoming light flux was measured carefully at the sample position before and after each photocatalytic activity test with a photo-radiometer (HD2302.0, Delta

Ohm/Italy) using a PAR VIS probe (400–700 nm). Typical VIS-light photon flux used in the current experiments was within 450–500 μ mol/ (m²sec). Reactor temperature remained within 23–42 °C during a typical 60 min photocatalytic activity test with a VIS light source. In each performance analysis test, 200 mg of photocatalyst was packed in a 2 mm \times 40 mm \times 40 mm poly methyl methacrylate (PMMA) sample holder and placed into the flow reactor. In order to quantify the photocatalytic activity and selectivity, at the end of a typical 1 h photocatalytic activity test, obtained raw data was processed in order to obtain various figures of merit (such as percent photonic efficiency towards NO $_{\rm x}$ storage, percent photonic efficiency for NO $_{\rm 2}$ generation/release, percent NO conversion, percent selectivity towards NO $_{\rm x}$ storage, and DeNO $_{\rm x}$ index). Details of these figures of merit are explained in detail in the Supporting Information (SI) section.

3. Results and discussion

3.1. Electron microscopy analysis via TEM and STEM

Surface morphology of as-synthesized Fe₃O₄/mpg-CN500 sample was investigated via high-resolution TEM, STEM and High Angle Annular Dark Field (HAADF) imaging techniques as shown in Fig. 1. Since Fe₃O₄/mpg-CN500 sample revealed the highest photocatalytic performance (vide infra), this particular catalyst was chosen for the electron microscopy analysis. Images in Fig. 1a and b provide multiple views of the 2D (sheet-like) structure of the mpg-CN500 support material containing thin layers of mpg-C₃N₄ in Fe₃O₄/mpg-CN500 sample. Fig. 1c presents HAADF-STEM images revealing a general view of the dispersion and particle size variation of the Fe₃O₄ nanoparticles on the CN500 support. It is seen in Fig. 1c that average particle size of the Fe₃O₄ nanoparticles on the mpg-CN500 support material ranges within ca. 8-11 nm. Fig. 1d shows a HR-TEM image of a Fe₃O₄ nanoparticle with a diameter of ca. 8 nm located on the edge of the mpg-CN500 (i.e. mpg-C₃N₄) support material. Lattice fringes measured on this Fe₃O₄ nanoparticle yields an average spacing of 0.254 nm, which is in accordance with the (311) plane of magnetite (JCPDS card no. 19-0629). It should be noted that g-C₃N₄ has a distinct fringe spacing of 0.325 nm that corresponds to the inter-layer structural packing of (002) crystal plane (JCPDS 87-1526) [22,28].

3.2. Crystal structure analysis via XRD

Fig. 2 presents the XRD patterns of the synthesized mpg-CN450, mpg-CN500 and mpg-CN550 samples as well as the Fe₃O₄/mpg-CN500 catalyst along with the pure Fe₃O₄ reference material (JCPDS Card No: 19-0629). For pure mpg-CN450, mpg-CN500 and mpg-CN550 samples, two main diffraction peaks are observed at 27.5° and 13.2° which confirm the formation of graphitic carbon nitride (JCPDS Card No: 87-1526). Strong diffraction peak at 27.5° is a characteristic interplanar stacking feature of the conjugated aromatic system, indexed for mpg-C₃N₄ as the (002) peak. The small diffraction peak at around 13.2° corresponds to intralayer/in-plane diffraction of the continuous tri-s-triazine units and indexed as (100) [29]. With increasing calcination temperature within 450–550 °C, diffraction signal located at $2\theta = 27.5^{\circ}$ became sharper and the intensity of the diffraction peak at $2\theta = 13.2^{\circ}$ increased monotonically. Furthermore, weak features located at $2\theta = 11.1^{\circ}$ and 17.6° (due to impurities and minority phases) disappeared. These observations are in line with the ordering of the crystal structure of the mpg-C₃N₄ support material. Anchoring of the Fe₃O₄ NPs on mpg-C₃N₄ resulted in the appearance of a new set of characteristic diffraction signals that are in very good agreement with that of pristine Fe₃O₄ (JCPDS Card No: 19-0629) [22], which is also shown in Fig. 2. Addition of Fe₃O₄ NPs also led to broadening of the main mpg-C₃N₄ diffraction feature at $2\theta = 27.5^{\circ}$ indicating decreased crystallinity and increased disorder in the carbon nitride domains due to the presence of Fe₃O₄ NPs and host-guest interactions [30]. This observation may also imply a

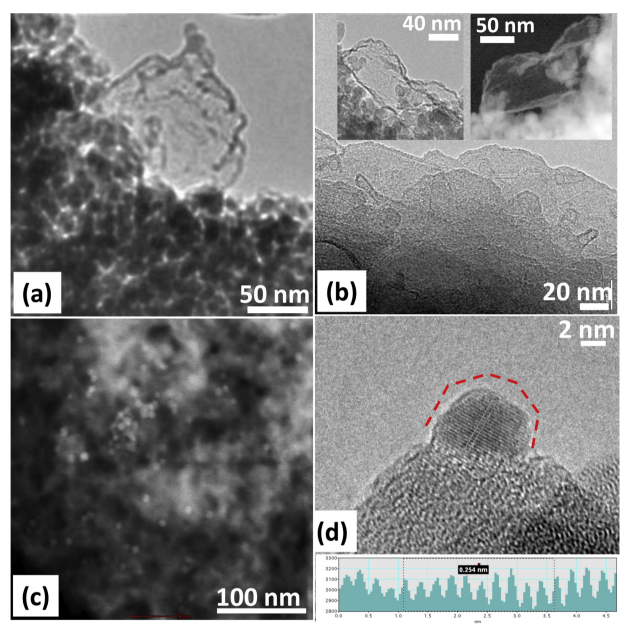


Fig. 1. (a–b) Multiple views of the 2D (sheet-like) structure of the mpg-C₃N₄ support material in Fe₃O₄/mpg-CN500. (c) HAADF-STEM image of Fe₃O₄/mpg-CN500 sample. (d) High-resolution TEM image of a Fe₃O₄ nanoparticle on the Fe₃O₄/mpg-CN500 photocatalyst (inset shows the distances between lattice fringes measured along the dashed white lines).

limited extend of incorporation of Fe species into the $\it mpg\text{-}C_3N_4$ matrix/layers.

3.3. Investigation of the surface composition via XPS

XPS measurements were performed in order to investigate the surface chemistry and functional groups present on the synthesized $\it mpg-G_3N_4-$ based photocatalysts. Fig. 3a-d show the C1s, N1s, Fe2p and O1s regions of the XP spectra corresponding to the Fe₃O₄/mpg-CN500 sample, respectively. In one of our former reports [17], we have provided a detailed discussion about the interpretation of the XP spectra of mpg-C₃N₄ materials that have been synthesized in a similar (though not identical) fashion in the absence of Fe₃O₄. Thus, in the light of the information provided in this aforementioned report [17], we can discuss the prominent XPS features of the currently synthesized Fe₃O₄/mpg-CN500 photocatalyst.

Fig. 3a presents the C1s region of the XP spectra of Fe₃O₄/mpg-

CN500 with a complex and a convoluted structure. It exhibits an intense high binding energy (B.E.) peak at 288.4 eV, which can be assigned to sp²-hybridized carbon species. The feature at 286.9 eV can be attributed to a variety of carbon species bound to N-containing species such as, spbonded carbon atoms in C≡N groups [31], sp²-hybridized s-triazine aromatic ring carbon, $(-N-)_2-C = N$ or sp^2 -hybridized s-triazine aromatic ring carbon coordinated to the terminal amino group, N-C(-NH₂) = N or carbon species bound to O-containing species. Note that readily detectable amounts of oxygen species are commonly observed in mpg-C₃N₄ structures, which can be located at the termination sites or defects/impurities in the s-triazine aromatic ring system of mpg-C₃N₄ [17]. C1s feature at 284.6 eV in Fig. 3a can be attributed to the C species that are mostly bound to other carbon atoms and also to adventitious carbon. Since in an ideal mpg-C3N4 structure containing interconnected s-triazine rings, there should not be any C-C/C = C linkages, presence of this feature implies the existence of carbonaceous impurities (e.g. graphite, graphene, graphene oxide etc.) in the mpg-

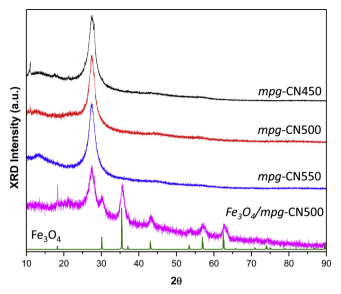


Fig. 2. XRD profiles of mpg-CN450, mpg-CN500, mpg-CN550 and Fe_3O_4/mpg -CN500 photocatalysts and pure Fe_3O_4 reference material.

 C_3N_4 system. This was also observed in our former report on similar *mpg*- C_3N_4 materials [17,32]. Finally, XPS spectrum in Fig. 3a also contains a broad low B.E feature (281.6 eV), which can be tentatively assigned to C species connected to the Fe species of the Fe₃O₄ NPs and C species connected to the residual Si species originating from the colloidal silica template used in the synthesis [33,34].

N1s region of the XP spectra of Fe₃O₄/mpg-CN500 can be seen in Fig. 3b. The broad and convoluted N1s feature located at 401.9 eV can be ascribed to N connected to O species at terminations or defects [17,35] and/or C-NH₂ (amino) functionalities with a possible contribution from differential charging effects [36]. The N1s signal present at 399.0 eV can be assigned to C-N-C species in s-triazine rings [17]. The most prominent N1s feature observed at 397.2 eV is associated with N atoms connected to C atoms in the s-triazine rings and/or pyridinic Nspecies [37]. Remaining minor peak at 395.1 eV in Fig. 3b with a low B.E. is tentatively assigned to N bound to Fe species (e.g. Fe-NH₂, Fe-N etc.) [38]. It should be noted that XPS analysis of semiconductors such as graphitic carbon nitride requires special attention as the XP spectra is extremely sensitive to the charge compensation (i.e. electron flood gun parameters) that is used during the XPS analysis. Along these lines, referencing of the B.E. positions are also difficult to accomplish since: a) due to the large C-content of the sample, it is not perfectly clear which C1s peak corresponds to the adventitious C; b) since the sample is not electrically conducting and also not homogeneous in terms of its surface composition/surface electronic structure, differential charging phenomena can severely shift, broaden, intensify or attenuate C1s signals in a drastic manner. Example XPS measurements with different chargecompensation parameters are presented in the Supporting Information section (Fig. S4). Nevertheless, general features of the currently presented C1s and N1s spectra are in line with our former studies [17].

Fe2p B.E. region of the XP spectra of Fe₃O₄/mpg-CN500 are presented in Fig. 3c. It is known that Fe₃O₄ (magnetite) typically reveals Fe2p_{1/2} and Fe2p_{3/2} XPS features at 724.1 \pm 1 eV and 710.6 \pm 1 eV, while these features characteristically appear at 724.6 \pm 1 eV and 711.0 \pm 1 eV for Fe₂O₃ (hematite), respectively. Furthermore, another

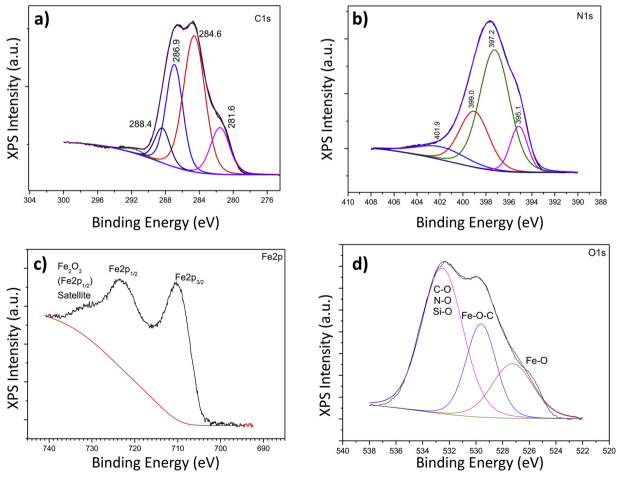


Fig. 3. XPS data for (a) C1s, (b) N1s, (c) Fe2p, and (d) O1s regions of Fe₃O₄/mpg-CN500 photocatalyst.

characteristic identifier for Fe_2O_3 is the satellite pair observed at 729.5 and $718.8\,eV.$ On the other hand, Fe_3O_4 does not reveal such satellites [39,40]. Along these lines, XPS data in Fig. 3c suggest that both Fe_3O_4 and Fe_2O_3 species coexist in the $Fe_3O_4/mpg\text{-CN}500$ system. Lack of any Fe_2O_3 diffraction features in the XRD data presented in Fig. 2 indicates that Fe_2O_3 species may exist as disordered/amorphous minority domains or they may exist as an amorphous surface oxide covering the Fe_3O_4 nanoparticles forming a core-shell structure. The latter structure can be formed upon oxidation of Fe_3O_4 nanoparticles during calcination in air to form a surface Fe_2O_3 overlayer.

O1s XP spectrum for Fe₃O₄/mpg-CN500 given in Fig. 3d reveals a complex and a convoluted structure possessing contributions from a large variety of oxygen species. Hence, we will not attempt a detailed deconvolution of this data but rather mention some of the possible prominent species that might be associated with the spectrum. O1s data in Fig. 3d shows features at 532.5 eV, 529.6 eV and 527.2 eV. The feature observed at 532.5 eV is most likely associated with oxygen species bound to C, N and Si in different configurations [35,41,42]. The O1s feature located at ca. 529.6 eV can be attributed to Fe-O-C species while the shoulder at 527.2 eV is assigned to Fe-O species and/or differential charging features [41,43].

3.4. Bulk chemical composition via ATR-FTIR spectroscopy

ATR-FTIR spectroscopy was used to investigate the vibrational characteristics and surface functional groups of the synthesized mpg- C_3N_4 photocatalysts (Fig. 4). The broad bands in the 3000–3500 cm⁻¹ region can be attributed to the N-H stretchings corresponding to the hydrogenation of terminal nitrogen atoms in mpg-C₃N₄ structures or N-H functionalities located at the surface defect sites [29]. The relatively weaker band at 2181 cm⁻¹ can be assigned to the $-C \equiv N$ triple bond stretchings of cyano groups in the mpg-C₃N₄ structure. These cyano/cyanide as well as other uncondensed defects like NH2/NH, -OH, -COOH are unavoidable as mpg-C₃N₄ structure evolves through the formation of several intermediates, such as cyanuric acid and melon [17]. It is interesting to note that the peak at 2181 cm⁻¹ is more prominently observed only at higher processing temperature i.e. $550~^{\circ}$ C or by the introduction of Fe₃O₄ NPs. IR adsorption band at 1617 cm⁻¹ can be ascribed to the C=N stretching vibration modes, while the 1568, 1408, 1317 and 1238 cm⁻¹ vibrational features are attributed to the typical stretching modes of the C-N aromatic heterocycles [44]. The absorption band at 806 cm⁻¹ is assigned to the out of plane bending

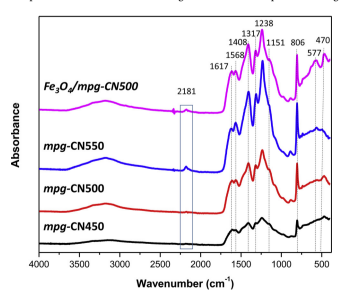


Fig. 4. ATR-FTIR spectra of mpg-CN450, mpg-CN500, mpg-CN550 and ${\rm Fe_3O_4/mpg\text{-}CN500}$ photocatalysts.

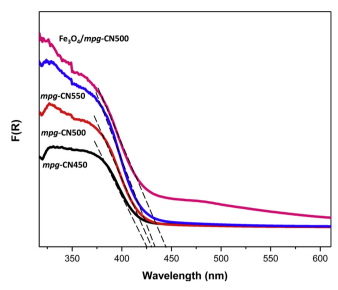


Fig. 5. a) DR-UV-VIS spectra of *mpg*-CN450, *mpg*-CN500, *mpg*-CN550 and Fe_3O_4/mpg -CN500 photocatalysts; b) Schematic representation of one of the possible Fe inclusion sites inside *mpg*-CN500 layers.

mode of triazine units [29]. The bands of the mpg-C₃N₄ photocatalysts become sharper with increasing calcination temperature because of the more ordered packing of the polymeric triazine units. The FTIR spectrum of Fe₃O₄/mpg-CN500 in Fig. 4 also shows two additional low frequency stretching bands that can be attributed to Fe₃O₄. The band at 577 cm⁻¹ is associated with the Fe-O deformation in octahedral and tetrahedral sites, while the band at 470 cm⁻¹ can be assigned to Fe-O deformation in octahedral sites of Fe₃O₄ [45,46]. In addition, 1148, 975 and 885 cm⁻¹ bands can be ascribed to residual ethanol which was used for washing purpose at the end of synthesis protocol [47,48].

3.5. Electronic band gap analysis via DR-UV-VIS spectroscopy

DR-UV-VIS spectra of the synthesized pure and Fe-doped mpg-C₃N₄ photocatalysts are shown in Fig. 5. All of the mpg-C₃N₄ photocatalysts show comparable absorption line shape due to the excitation of electrons and charge transfer from valence band comprised mainly of N2p states to the conduction band which is mostly formed by C2p states [49]. Increasing the calcination temperature from 450 °C to 550 °C during the mpg-C₃N₄ synthesis protocol leads to a gradual red shift in the absorption edge which can be associated with the enhancement of electron delocalization and conjugation in the aromatic sheets as a result of increasing crystallographic order. Subsequently, the band gap energy decreases from ca. 2.74 eV to 2.72, and 2.70 eV, as the calcination temperature is gradually increased from 450 °C to 500 °C, and 550 °C, respectively. Incorporation of Fe₃O₄ into mpg-CN500 structure has two noticeable effects in the DR-UV-VIS spectra. Firstly, it enhances the absorption intensity. Secondly, it leads to a further red shift in the absorption edge (i.e. 2.65 eV). This additional red shift in the absorption edge of Fe₃O₄/mpg-CN500 photocatalyst may facilitate the electronhole pair generation under VIS-light irradiation, which in turn, can result in improved photocatalytic efficiency [50]. Fe₃O₄ NPs are dispersed on/within the layers and pores of mpg-C₃N₄. We believe that -CN, -NH₂, -NO_x, -C_xH_vO_z terminations/functional groups/structural defects of mpg-C₃N₄ provide binding sites for Fe₃O₄ NPs. In addition, oleylamine capping agent of Fe₃O₄ NP might also provide additional binding capabilities with the mpg-C₃N₄ layers. In addition to these, other point defects, extended defects (e.g. edges, kinks), corners and/or termination sites could also be other possible sites for Fe₃O₄ NP accommodation.

3.6. Specific surface area analysis via BET

Specific surface area (SSA) of mpg- C_3N_4 materials were determined by BET nitrogen adsorption isotherms. The specific surface area of mpg-CN450, mpg-CN500, mpg-CN550 and Fe_3O_4/mpg -CN500 photocatalysts were found to be 67, 140, 165 and 130 m^2/g , respectively. The increase in SSA with increasing calcination temperatures is due to the enhanced endothermic decomposition of guanidine hydrochloride at higher temperatures resulting in the formation of a porous mpg- C_3N_4 network with a higher SSA. On the other hand, anchoring of Fe_3O_4 NPs on mpg- C_3N_4 leads to a minor decrease in SSA, which can be attributed to the partial blocking of the mpg- C_3N_4 pores by Fe_3O_4 NPs with an average diameter of ca. 10 nm (Fig. 1a andb).

3.7. Photocatalytic $NO_x(g)$ oxidation and storage (PHONOS) in solid state

In order to demonstrate catalytic activity of the currently synthesized mpg-C₃N₄ photocatalysts, we performed photocatalytic NO_x(g) oxidation and storage (PHONOS) tests using a custom-made photocatalytic flow reactor. It is well known that the ultimate predominant form of the stored NO_x species on the photocatalyst surface is nitrate and HNO₃ species [51]. The term "NO_x Storage" is a very commonly used term in catalysis, particularly for DeNOx catalytic converters used in tail-pipe emission control systems of diesel engines. Such non-photocatalytic DeNOx catalysts (e.g. NOx Oxidation and Storage, NSR catalysts, also called Lean NOx Traps, LNT) are also commercially utilized, where NO(g) and NO₂(g) are stored on solid oxide catalysts in the form of nitrites and nitrates [6-8]. Along these lines, VIS-light induced removal of NO(g) was monitored under in-situ conditions for different mpg-C₃N₄ photocatalysts. Fig. 6 shows a typical set of time-dependent NO(g), NO₂(g) and total NO_x(g) concentration profiles as a function of irradiation time during the NO photo-oxidation over Fe₃O₄/mpg-CN500 photocatalyst. In the first stage of the photocatalytic activity tests, a synthetic polluted air gas mixture containing ca. 1 ppm NO(g) was fed to the photocatalyst surface under dark conditions. During this initial phase (i.e. first 15 min), a minor transitory fall in the total NO_x(g) and NO(g) concentrations was observed due to adsorption of NO_x species on the reactor lines, expansion of the gas in the reactor as well as non-

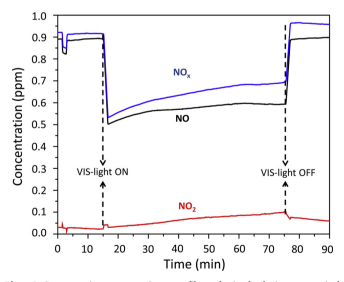


Fig. 6. Concentration *versus* time profiles obtained during a typical Photocatalytic $NO_x(g)$ Oxidation and Storage (PHONOS) activity test performed on Fe_3O_4/mpg -CN500 photocatalyst in a custom-made photocatalytic flow reactor. Red, black and blue traces correspond to $NO_2(g)$, NO(g) and total NO_x (*i.e.* $NO(g) + NO_2$) concentrations measured (with a 1 Hz acquisition rate) as a function of time during the photocatalytic activity test. Feed composition: $N_2(g)$ 0.750 SLM, $O_2(g)$ 0.250 SLM, and NO(g) 0.010 SLM (100 ppm NO (g) diluted in balance $N_2(g)$, RH 50% at 23 °C.

photocatalytic adsorption of NO_x on the photocatalyst surface. In addition, a tiny amount of $NO_2(g)$ was produced due to thermal catalytic disproportionation processes occurring on the catalyst surface. Following the saturation of the reactor system and photocatalyst surface, $NO_x(g)$ and NO(g) levels quickly returned to the original inlet concentration and reached a steady state in dark conditions.

Next, VIS-light irradiation was turned on after the first ca. 15 min (Fig. 6) and a drastic fall in the NO(g) and total NO_x(g) concentrations was detected along with a small increase in the NO₂(g) level. While the latter observation suggests the photocatalytic oxidation of NO(g) into NO₂(g), fall in the NO(g) and total NO_x(g) concentrations indicates the solid state storage of NO(g) and NO₂(g) in the form of chemisorbed NO₂, nitrites and/or nitrates on the mpg-C₂N₄ surface [8,52]. In principle, N2(g) and/or N2O(g) can also be produced as a result of direct photocatalytic decomposition and photo-reduction of NO(g) [53]. However, this is known to be a relatively inefficient reaction pathway, particularly in the presence of H₂O(g) and thus can readily be ruled out in the current study as a minor photocatalytic route [54]. It is apparent in Fig. 6 that photocatalytic NO_x abatement action continues in an uninterrupted manner after this initial stage during the entire duration of the activity test. Thus, total NOx abatement effect can be calculated by integrating the relevant traces for the total PHONOS test duration (see SI section for the details of such calculations).

Photocatalytic NO_x(g) oxidation and storage (PHONOS) activity tests were performed for all of the mpg-C₃N₄-based photocatalysts and compared with that of a commercial benchmark titania P25 photocatalyst under identical experimental conditions (Figs. 7-9). Fig. 7 presents the photocatalytic activity data in terms of NO_x storage photonic efficiency % and NO2(g) production photonic efficiency % values (which are normalized by the corresponding VIS-photon flux values), while Fig. 8 illustrates the absolute NO(g) conversion % and NO2 storage selectivity % values. Meanwhile, overall photocatalytic NO_x abatement performances of all of the investigated photocatalysts are presented in Fig. 9 in terms of their corresponding DeNO_x index values. Occupational Safety and Health Administration (OSHA), Association Advancing Occupational and Environmental Health (ACGIH) and The National Institute for Occupational Safety and Health (NIOSH) regulations suggest an environmental hazard limit value of 25 ppm for NO(g). On the other hand, corresponding limit value for NO₂(g) varies from 1 to 3 ppm, making NO₂(g) 8–25 times more toxic than NO(g) [55]. Based on these references, we have conservatively assigned a relative toxicity value of 1 to NO(g) and 3 to NO₂(g) [15] (see SI for more info). It is worth mentioning that per cent photonic efficiencies presented in Fig. 7 may seem relatively low (i.e. < 0.03%) and this is due to the VIS photon flux normalization. On the other hand, it should be noted that the corresponding percent conversion and selectivity values obtained

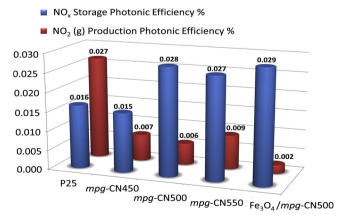


Fig. 7. NO_x storage photonic efficiency % (blue bars) and NO_2 (g) production photonic efficiency % (red bars) values for mpg-CN450, mpg-CN500, mpg-CN500 and Fe_3O_4/mpg -CN500 photocatalysts obtained via Vis-light irradiation.

NO Conversion %

■ NO, Storage Selectivity %

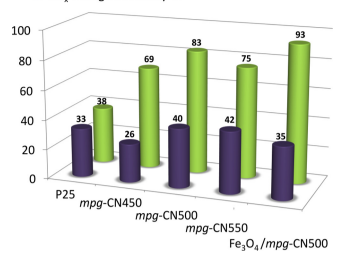


Fig. 8. NO(g) conversion % (purple bars) and NO $_2$ storage selectivity % (green bars) values for mpg-CN450, mpg-CN500, mpg-CN550 and Fe $_3$ O $_4$ /mpg-CN500 photocatalysts obtained via Vis-light irradiation.

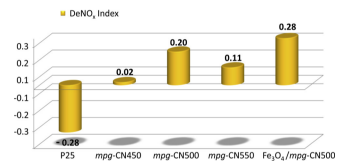


Fig. 9. $DeNO_x$ index values for mpg-CN450, mpg-CN500, mpg-CN550 and Fe_3O_4/mpg -CN500 photocatalysts obtained via Vis-light irradiation.

without photon flux normalization (Fig. 8) are quite high (i.e. 33-93 %)

It is important to note that absolute photocatalytic conversion and selectivity values are highly sensitive to reaction conditions and reactor design. This has been an important and well-known challenge for the comparison of literature data obtained by different groups using different experimental parameters. Hence, use of a benchmark catalyst and reporting "relative values with respect to a benchmark" is the most effective method for data comparison. On the other hand, choice of the benchmark catalyst should be made in such a way that it allows comparison of the maximum number of independent research studies published in the literature. Along these lines, P25 form of titania was chosen in the current work as the benchmark catalyst since this is probably the most commonly used photocatalyst in the literature.

It is clear from Figs. 7 and 8 that titania P25 commercial benchmark photocatalyst has a reasonably high NO_x storage efficiency. On the other hand, this titania benchmark catalyst simultaneously produces a very large quantity of unwanted $NO_2(g)$. Despite the fact that the adsorption capacity of TiO_2 for $NO_2(g)$ is much higher than that for NO(g) [56], photocatalytic $NO_2(g)$ production rate and the total amount of NO_2 generated readily overwhelms the NO_x adsorption capacity of titania leading to unwanted $NO_2(g)$ slip/release into the atmosphere. As $NO_2(g)$ is a much more toxic pollutant than NO(g), P25 does not qualify as an efficient photocatalyst for NO_x abatement under visible light irradiation, evident by the extremely negative $DeNO_x$ index of P25 (-0.28) given in Fig. 9.

When the corresponding photocatalytic performances of $mpg\text{-}C_3N_4$ -based photocatalysts are investigated (Figs. 7–9), striking improvements can be readily noticed as opposed to that of titania P25 commercial benchmark photocatalyst such as increased percent NO_x storage photonic efficiency and decreased $NO_2(g)$ production photonic efficiency (Fig. 7), increased NO conversion %, increased NO_x storage selectivity % (Fig. 8), and increased $DeNO_x$ index (Fig. 9) values.

Comparison of the relative photocatalytic performance of pure $\it mpg-C_3N_4$ photocatalysts (*i.e.* $\it mpg-CN450$, $\it mpg-CN500$, $\it mpg-CN550$) in Figs. 7–9 clearly reveals a volcano-plot like behavior suggesting that in the absence of Fe incorporation, $\it mpg-CN500$ photocatalyst demonstrates the optimum performance by maximizing photocatalytic NO_x oxidation and storage, while minimizing the NO₂(g) release to the atmosphere. As opposed to P25 titania commercial benchmark photocatalyst, NO conversion % was improved from 33% to 40% and NO_x storage selectivity % was enhanced from 38% to 83% on $\it mpg-CN500$ (Fig. 8). On the other hand, negative DeNO_x index of P25 titania commercial benchmark photocatalyst (-0.28) was radically surpassed by the $\it mpg-CN500$ photocatalyst (+0.20, Fig. 9). In other words, while P25 titania commercial benchmark photocatalyst does not reveal a net NO_x abatement effect under VIS irradiation, $\it mpg-CN500$ photocatalyst reveals a strong photocatalytic detoxification action under VIS-light.

Superior photocatalytic performance of mpg-CN500 over titania P25 can originate from two major functional improvements associated with mpg-CN500. Firstly, mpg-CN500 can perform NO(g) photo-oxidation with a greater effectiveness, possibly due to its favorably lower electronic band gap (2.72 vs. 3.1 eV), greater electron-hole generation capability, better charge migration, longer life-time of charge carriers and well-dispersed photocatalytic active sites over the mesoporous mpg-C₃N₄ 2D-layers allowing efficient harvesting of VIS-light. Secondly, surface functional groups of mpg-CN500, structural defects and the unique termination sites as well as the relatively higher specific surface area of mpg-C₃N₄ as compared to that of P25 results in enhanced NO_xcapture and storage in the solid state on mpg-CN500. It is worth mentioning that enhanced photocatalytic performance of mpg-CN500 over P25 titania or other mpg-C₃N₄ photocatalysts cannot be solely attributed to the greater SSA of the former photocatalyst but rather involves various structural/electronic/surface chemical properties of mpg-CN500 discussed above and further examined in the later part of this report. Different photocatalysts can exhibit same photoactivity despite having comparatively different surface areas, because activity is ratelimited by photoreaction rather than adsorption [57]. For instance, although SSA of mpg-CN550 (165 m²/g) is greater than that of mpg-CN500 (140 m²/g), mpg-CN550 demonstrates a lower photocatalytic activity as compared to mpg-CN500.

In their detailed mechanistic study, Bloh et al. [51] suggested that there may be two opposing molecular pathways governing the overall selectivity in PHONOS process. It is well known that the ultimate predominant form of the stored NO_x species in the solid state is nitrate and HNO₃ species adsorbed on the photocatalyst surface. Thus, in order to maximize the activity and selectivity of the photocatalyst, one needs to maximize nitrate coverage on the surface without facilitating NO₂(g) release. However, it was reported that [51], there is a trade-off between nitrate coverage and NO₂(g) release, where increasing nitrate surface coverage may trigger unwanted NO₂(g) production with the assistance of conduction band electrons generated by the photon absorption. In other words, it was proposed that the origin of NO2(g) generation in PHONOS process is the reduction of surface nitrates. An effective strategy to circumvent this problem was suggested to consider electron scavengers which can suppress nitrate reduction by facilitating O2 reduction to form surface superoxide and hydroperoxyl radicals (with the assistance of H⁺(ads) species). Therefore, the use of co-catalysts which can scavenge electrons from the nitrate reduction pathways, while facilitating molecular oxygen reduction could significantly enhance selectivity of the PHONOS systems without sacrificing their photocatalytic activity (i.e. conversion). Along these lines, FeOx

semiconductor nanoparticles containing $\mathrm{Fe^{2^+}}$ and $\mathrm{Fe^{3^+}}$ redox sites (such as the currently used $\mathrm{Fe_3O_4}$ nanoparticles containing minority $\mathrm{Fe_2O_3}$ surface domains) could be a promising alternative to boost the selectivity of $\mathit{mpg\text{-}C_3N_4}$ photocatalysts without sacrificing their photocatalytic activity.

As stated above, NO2 is a poisonous intermediate and must be avoided. Adsorbed oxygen on the catalyst surface can compete with nitrates for electrons and suppress the formation of NO2. However, the reduction of nitrates to NO2 is thermodynamically favored as compared to molecular oxygen reduction. To address this problem, an FeOx cocatalyst can be introduced which can promote oxygen reduction. Until now, high selectivity towards solid state NO_x storage and prevention of gaseous NO₂ formation have been achieved either via utilization of precious metals [58-60] or this selectivity increase was achieved at the expense of a sacrifice in absolute activity [15,61]. In this regard, nonprecious transition metals present an effective and convenient alternative to counter these drawbacks and increase the oxygen reduction capabilities of photocatalysts [62,63]. Here, we propose the introduction of Fe₃O₄ onto mpg-C₃N₄ photocatalyst to suppress the release of NO₂ during photo-oxidation of NO. The acceleration in the oxygen reduction with the assistance of Fe₃O₄ can allow a larger number of holes to oxidize the desired compounds (i.e. NO and NO₂) [64]. The resultant modification can improve the optical absorption range, as well as the charge carrier transfer rate in Fe₃O₄/mpg-C₃N₄, ultimately resulting in improved activity and selectivity [65].

Figs. 7–9 clearly demonstrate that this is an efficient strategy. It is evident that incorporation of FeO_x species into the optimized $mpg\text{-}C_3N_4$ photocatalyst (i.e. mpg-CN500) to form $Fe_3O_4/mpg\text{-}CN500$ photocatalyst leads to a further enhancement of overall photocatalytic performance by leading to a ca. 40% boost in $DeNO_x$ index (Fig. 9) from +0.20 (mpg-CN500) to +0.28 ($Fe_3O_4/mpg\text{-}CN500$) under VIS-light illumination. This is accomplished by a substantial increase in NO_x storage selectivity to 93% with only a minor decrease in NO_x conversion % to 35% (Fig. 8).

Currently observed synergistic effect associated with the FeO_x doping of $mpg\text{-}C_3N_4$ photocatalysts can also be attributed to the lowering of the electronic band gap energy (2.65 eV) and reduced recombination rate of photo-generated electron-hole pairs. The latter argument can be supported by considering the fact that the reduction potential of Fe^{2+}/Fe^{3+} is below the conduction band of g-C₃N₄ [30]. We believe that Fe^{2+}/Fe^{3+} species can act as electron scavengers in $Fe_3O_4/mpg\text{-}CN500$ photocatalysts where photo-generated electrons are trapped by the Fe_3O_4 doping sites, resulting in a decrease in the recombination rates of photo-generated electron–hole pairs.

As can be seen in Fig. 8, NO conversion % is slightly lower for Fe_3O_4/mpg -CN500 (35%) as compared to that of mpg-CN500 (40%) which is probably due to the loss of specific surface area after doping (130 vs. 140 m^2/g , respectively). Fe_3O_4/mpg -CN500 photocatalyst performed remarkably better, when compared with similar g-C $_3N_4$ based photocatalyst systems in terms of conversion%, selectivity% and DeNOx Index under VIS light illumination [66–68]. Further studies are underway in our group towards exploring the optimum Fe_3O_4 loading.

3.8. Photoluminescence (PL)

In order to elucidate the origins of the differences in photocatalytic NOx oxidation performance of the currently investigated samples, PL experiments were carried out. PL experiments revealed information regarding the migration, transfer, and recombination processes of photoinduced electron – hole pairs in the *mpg*-C₃N₄ systems. Fig. 10 shows the PL spectra of *mpg*-CN450, *mpg*-CN500, *mpg*-CN550 and Fe₃O₄/*mpg*-CN500 obtained with an excitation wavelength of 390 nm at room temperature. A strong emission peak was observed for *mpg*-CN450 indicating higher recombination rate of excited electrons and holes, ultimately leading to a lower photocatalytic efficiency [69]. As the *mpg*-C₃N₄ synthesis temperature increases, PL emission peak intensity

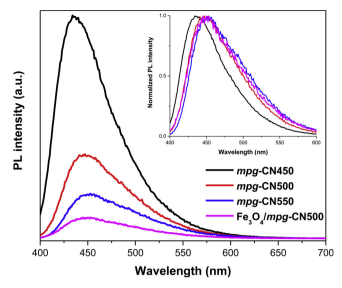


Fig. 10. PL spectra of mpg-CN450, mpg-CN500, mpg-CN550 and Fe $_3$ O $_4$ /mpg-CN500 photocatalysts ($\lambda_{\rm ex}=390$ nm). Intensity-normalized spectra are shown in the inset.

decreases significantly for mpg-CN500 and mpg-CN550, resulting in better photocatalytic performance of these two materials as compared to mpg-CN450. For Fe₃O₄/mpg-CN500, PL intensity further decreases in a notable manner, indicating a much lower recombination rate of photoinduced electron - hole pairs. This probably happens because photoinduced electrons can easily migrate from the conduction band (CB) of mpg-CN500 to the CB of Fe₃O₄ (i.e. away from the holes), thanks to the energy matching band structure of the Fe₃O₄/mpg-CN500 composite system [49]. Moreover, Fe₃O₄ has a relatively fast electron transport rate due to its high conductivity which suppresses the direct recombination of photo-induced electron - hole pairs in the Fe₃O₄/ mpg-CN500 system [22]. The electrons in the CB of Fe₃O₄ are good reductants that can efficiently reduce the O2 species adsorbed on the photocatalyst surface to various reactive species (e.g. superoxide and hydroperoxyl radicals), consequently suppressing the reduction of nitrates to NO2 and leading to enhanced photocatalytic oxidation of NOx.

It is interesting to note here that, although PL spectra show significantly low emission intensity for *mpg*-CN550, it has comparatively low photocatalytic efficiency as compared to *mpg*-CN500 (Figs. 7–9). Upon closer inspection of IR spectra in Fig. 4, it becomes evident that *mpg*-CN550 structure has cyanide/cyano defects (2181 cm $^{-1}$) which could reduce the electron mobility on the π conjugation plane of g-C₃N₄. For *mpg*-CN550, we believe that non-radiative (thermal) relaxation channels could be responsible for the decrease in the free charge carrier concentration, subsequently resulting in reduced photocatalytic ability [70,71]. To a lesser extent, this adverse effect can also be observed for the NO conversion on Fe₃O₄/*mpg*-CN500 photocatalyst (Fig. 4 and 8).

The inset in Fig. 10 shows that centers of PL spectra shift with the increase of processing temperatures (435 nm for *mpg*-CN450, 447 nm for *mpg*-CN500 and 453 nm for *mpg*-CN550), indicating the gradual development of defects in the material. Additionally, the full width at half maximum (FWHM) also becomes broader with the increasing temperature due to the changes in degree of thermal condensation and microstructural order in the *mpg*-C₃N₄ photocatalysts [72,73]. These results suggest that extent of polymerization of the precursor and the concentration of cyano/cyanide trap sites play an important role in the overall photocatalytic activity of the *mpg*-C₃N₄ photocatalysts. In addition, different precursors and utilization of various purification methods might also influence the type and concentration of defects in the *mpg*-C₃N₄ system, which may allow further fine tuning of photocatalytic performance [70].

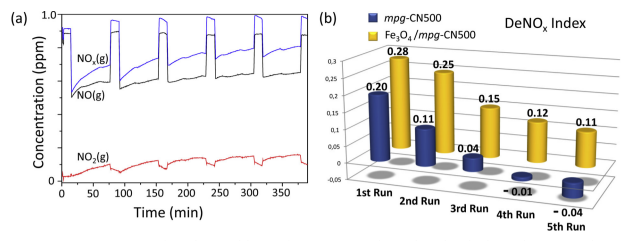


Fig. 11. (a) Concentration *versus* time curves for catalytic reusability test of Fe_3O_4/mpg -CN500 photocatalysts where the photocatalyst was exposed to 5 subsequent identical PHONOS runs under VIS-light irradiation, (b) corresponding $DeNO_x$ index values for mpg-CN500 and Fe_3O_4/mpg -CN500 photocatalysts calculated for each run of the reusability tests.

3.9. Photochemical stability of mpg-C₃N₄-based photocatalysts and reusability

Photochemical-stability and reusability of photocatalysts are crucial properties for end user applications. Thus, in order to demonstrate the reusability of the currently investigated materials, we performed a series of experiments. In these experiments, five successive VIS-light induced photo-oxidation tests were performed for two of the best performing photocatalyst (i.e. mpg-CN500 and Fe₃O₄/mpg-CN500) under identical reaction conditions. It must be stated here that no regeneration treatment was performed for these photocatalysts during these reaction cycles. As shown in Fig. 11, the DeNO_x index value gradually decreased for mpg-CN500 after each cycle and became negative after the fourth cycle. This monotonic attenuation in NOx abatement capability could be probably due to the continuous accumulation of nitrites on the photocatalyst surface diminishing the NO_x storage capacity and facilitating NO₂(g) generation by nitrate reduction. While a monotonic decrease in DeNO_x index was also visible for the Fe₃O₄/mpg-CN500 photocatalyst, attenuation in the DeNO_x index values was significantly smaller in magnitude resulting in positive DeNO_x index values for all cycles. Note that identical re-usability experiments performed with P25 titania benchmark catalysts (data not shown) also revealed a monotonically decreasing DeNO_x index values where the values ranged between -0.30 and -0.40.

These results clearly indicate that the Fe $_3$ O $_4$ /mpg-CN500 photocatalyst has excellent long-term stability and enhanced activity. The slight decrease in photocatalytic activity of Fe $_3$ O $_4$ /mpg-CN500 photocatalyst with time is primarily due to the accumulation of surface nitrate species which can be easily regenerated by washing the photocatalyst with water [74]. In other words, Fe $_3$ O $_4$ /mpg-CN500 photocatalyst architecture can be considered as a highly active, selective, stable and an affordable material that can be synthesized readily in large scales and applied in outdoor NO $_x$ abatement applications.

4. Conclusion

In the current work, highly efficient Fe $_3$ O $_4$ NPs anchored mpg-C $_3$ N $_4$ photocatalysts were synthesized by a facile and cost-effective synthetic protocol. Fe $_3$ O $_4$ /mpg-CN500 photocatalyst revealed significantly superior photocatalytic NO $_x$ abatement performance under visible light as compared to the commercial P25 titania benchmark photocatalyst. Fe species in the Fe $_3$ O $_4$ /mpg-CN500 photocatalyst structure existed as FeO $_x$ nanoparticles with an average diameter of ca. 10 nm on mpg-C $_3$ N $_4$ surface. Incorporation of the Fe species to the photocatalyst formulation enhanced the total light absorption in the visible region, decreased the

electronic band gap of mpg-C₃N₄ and also generated structural defects resulting in crystallographic disorder. Introduction of FeO_x species significantly increased the oxygen reduction capacity of mpg-CN500 resulting in lower NO2 production which ultimately resulted in enhanced selectivity in photocatalytic NO_x abatement. Fe₃O₄/mpg-CN500 photocatalyst showed high activity, selectivity and stability even after five successive experimental runs without any regeneration step. Enhanced photocatalytic efficiency could be mainly attributed to the unique mesoporous structure, high surface area, enhanced charge separation efficiency and prolonged life time of charge carriers. Further studies are ongoing in our research group for elucidating the detailed photocatalytic mechanism, influence of Fe loading, effect of temperature and influence of regeneration methodology. We believe that Fe₃O₄/mpg-CN500 presents itself as an example of a new family of affordable, active, scalable and environmentally friendly photocatalytic materials which has a significant potential in environmental remediation and solar energy conversion applications.

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.apcatb.2019.02.067.

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